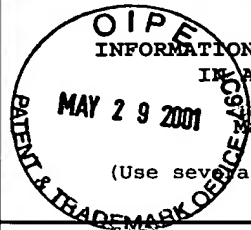
				ATTORNEY DOCKET NO. 3028.1000-000	APPLICATION NO. 09/590,211		
				APPLICANT Guy A. Rouleau and Bernard Brais			
				FILING DATE June 8, 2000	GROUP 1632		
U.S. PATENT DOCUMENTS							
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	AA						
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FOREIGN PATENT DOCUMENTS							
		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION YES NO
	AL	WO98/31800	23 JUL 98	PCT			
	AM	WO99/29896	17 JUN 99	PCT			
	AN						
	AO						
OTHER DOCUMENTS (Including Author, Title, Date, Pertinent Pages, Etc.)							
AR ✓	Akarsu, A.N., et al., "Genomic Structure of HOXD13 Gene: A Nine Polyalanine Duplication Causes Synpolydactyly in Two Unrelated Families," <i>Human Molecular Genetics</i> , 5(7): 945-952 (1996).						
AS ✓	Bienroth, S, et al., "Assembly of a Processive Messenger RNA Polyadenylation Complex," <i>The EMBO Journal</i> , 12(2): 585-594 (1993).						
AT ✓	Brais, B., et al., "Using the Full Power of Linkage Analysis in 11 French Canadian Families to Fine Map the Oculopharyngeal Muscular Dystrophy Gene," <i>Neuromuscular Disorder</i> 7(1):S70-S74 (1997).						
AU ✓	Brais, B, et al., "The Oculopharyngeal Muscular Dystrophy Locus Maps to the Region of the Cardiac α and β Myosin Heavy Chain Genes on Chromosome 14q11.2-q13," <i>Human Molecular Genetics</i> , 4(3): 429-434 (1995).						
AV ✓	Davies, S.W., "Formation of Neuronal Intranuclear Inclusions Underlies the Neurological Dysfunction in Mice Transgenic for the HD Mutation," <i>Cell</i> , 90:537-548 (1997).						
AW ✓	DiFiglia, M, et al., "Aggregation of Huntingtin in Neuronal Intranuclear Inclusions and Dystrophic Neurites in Brain," <i>Science</i> , 277: 1990-1993 (1997).						
AX ✓	Evans, G.A, et al., "High Efficiency Vectors for Cosmid Microcloning and Genomic Analysis," <i>Gene</i> , 79:9-20 (1989).						
AY ✓	Forood, B., et al., "Formation of an Extremely Stable Polyalanine β -Sheet Macromolecule," <i>Biochem. And Biophysical Res. Communications</i> , 211(1): 7-13 (1995).						
AZ ✓	Krause, S., et al., "Immunodetection of Poly(A) Binding Protein II in the Cell Nucleus," <i>Experimental Cell Res.</i> , 214: 75-82 (1994).						
EXAMINER				DATE CONSIDERED			

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<p>PTO-1449 REPRODUCED</p> <p>INFORMATION DISCLOSURE CITATION IN AN APPLICATION</p> <p>MAY 29 2001</p> <p>May 23, 2001</p> <p>(Use several sheets if necessary)</p>				ATTORNEY DOCKET NO. 3028.1000-000	APPLICATION NO. 09/590,211		
				APPLICANT Guy A. Rouleau and Bernard Brais			
				FILING DATE June 8, 2000	GROUP 1632		
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U.S. PATENT DOCUMENTS							
EXAM- INER INI- TIAL		DOCUMENT NUMBER	DATE	NAME	CLASS	SUB- CLASS	FILING DATE IF APPROPRIATE
FOREIGN PATENT DOCUMENTS							
		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION YES NO
OTHER DOCUMENTS (Including Author, Title, Date, Pertinent Pages, Etc.)							
AR2✓	Mundlos, S., et al., "Mutations Involving the Transcription Factor CBFA1 Cause Cleidocranial Dysplasia," <i>Cell</i> , 89: 773-779 (1997).						
AS2✓	Editorials, "DNA-Triplet Repeats and Neurologic Disease," <i>The New England Journal of Med.</i> , 335(16): 1222-1224 (1996).						
AT2✓	Scherzinger, E., et al., "Huntingtin-Encoded Polyglutamine Expansions Form Amyloid-Like Protein Aggregates In Vitro and In Vivo," <i>Cell</i> 90: 549-558 (1997).						
AU2✓	Tome, M.S., et al., "Nuclear Inclusions in Oculopharyngeal Dystrophy," <i>Act Neuropathol.</i> 49: 85-87 (1980).						
AV2✓	Muragaki, Y., et al., "Polyalanine Expansion in Synpolydactyly Might Result from Unequal Crossing-Over of HOXD13," <i>Science</i> 275: 406						
AW2✓	Wells, R.D., "Molecular Basis of Genetic Instability of Triplet Repeats," <i>The Journal of Biological Chem.</i> 271(6): 2875-2878 (1996).						
AX2✓	Wahle, E., et al., "Mammalian Poly(A)-Binding Protein II," <i>J. of Biological Chem.</i> , 268(4): 2937-2945 (1993).						
AY2✓	Wahle E., "A Novel Poly(A)-Binding Protein Acts As a Specificity Factor in the Second Phase of Messenger RNA Polyadenylation," <i>Cell</i> , 66: 759-768 (1991).						
AZ2✓	Nemeth, A., et al., "Isolation of Genomic and cDNA Clones Encoding Bovine Poly(A) Binding Protein II," <i>Nucleic Acids Res.</i> , 23(20): 4034-4041 (1995).						
AR3✓	Riggins, G.J., et al. "Human Genes Containing Polymorphic Trinucleotide Repeats," <i>Nat Genet</i> , 2(3):186-191 (1992).						
AS3✓	Brais, B., et al., "Short GCG Expansions in the PABP2 Gene Cause Oculopharyngeal Muscular Dystrophy," <i>Nature Genetics</i> 18: 164-167 (1998).						
EXAMINER				DATE CONSIDERED			

PTO-1449 REPRODUCED

ATTORNEY DOCKET NO.
3028.1000-000APPLICATION NO.
09/590,211APPLICANT
Guy A. Rouleau and Bernard BraisFILING DATE
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U. S. PATENT DOCUMENTS

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FOREIGN PATENT DOCUMENTS

		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION YES NO

OTHER DOCUMENTS (Including Author, Title, Date, Pertinent Pages, Etc.)

AT3 ✓	Lamartine, J. et al., "Cloning Sequencing and Chromosomal Assignment of a New cDNA Clone to Xq12-q13 and 14q11," EMBL Data Base Accession Number U12206 (1995).
AU3 ✓	"National Cancer Institute, Cancer Genome Anatomy Project (CGAP), Tumor Gene Index," EMBL Data Base Accession Number AA618589 (1997).
AV3 ✓	Sullivan, T.B.T., et al., "Oculopharyngeal Muscular Dystrophy (OPMD) - Report and Genetic Studies of an Australian Kindred," <i>Clinical Genetics</i> , 51: 52-55 (1997).
AW3 ✓	Bouchard, J.P. et al., "A Simple Test for the Detection of Dysphagia in Members of Families with Oculopharyngeal Muscular Dystrophy (OPMD)," <i>Can.J. Neurol. Sci.</i> 19(2):296-297 (1992).

EXAMINER

DATE CONSIDERED